

VEREYSKIY, N.G.; GANESHIN, G.S.; KRASNOV, I.I.; CHEMEKOV, Yu.F.

Fourth Congress of the International Association on Quaternary Research (INQUA). Sov.geol. 5 no.5:160-165 My '62. (MIRA 15:7)

1. Vsesoyuznyy nauchno-issledovatel skiy geologicheskiy institut i Vsesoyuznyy nauchno-issledovatel skiy institut gidrogeologii i inzhenernoy geologii.

(Geology, Stratigraphic--Congresses)

NALIVKIN, D.V., glav. red.; VERESHCHAGIN, V.N., zam. glav. red.;

MENNER, V.V., zam. glav. red.; CVECHKIN, N.K., zam. glav. red.[deceased]; SOKOLOV, B.S., zam. glav. red.; SHANTSER, Ye.V., zam. glav. red.; KELLER, B.M., otv. red. toma;

MODZALEVSKAYA, Ye.A., red.; CHUGAYEVA, M.N., red.;

GROSSGEYM, V.A., redaktor; KIPARISOVA, L.D., redaktor; KOROBKOV, M.A., red.; KRASNOV, I.I., red.; KRYMCOL'TS, T.Ya., red.; LIBROVICH, L.S., red.; LIKHAREV, B.K., red.; LUPPOV, N.P., red.; NIKIFOROVA, O.I., red.; QERUCHEV; S.V., red.; POLKANOV, A.A., red.[deceased]; RENGARTEN, V.P., red.; STEPANOV, D.L., red.; CHERNYSHEVA, N.Ye., red.; SHATSKIY, N.S., red. [deceased]; EBERZIN, A.G., red.; GOROKHOVA, T.A., red.izd-va; GUROVA, O.A., tekhn. red.

[Stratigraphy of the U.S.S.R. in fourteen volumes] Stratigrafiia SSSR v chetyrnadtsati tomakh. Moskva, Gosgeoltekhizdat. Vol.2. [Upper Pre-Cambrian] Verkhnii dokembrii. Otv. red. B.M. Keller. 1963. 716 p. (MIRA 17:1)

1. Chlen-korrespondent AN SSSR (for Sokolov).

NALIVKIN, V.D.; RONOV, A.B.; KHAIN, V.Ye.: VOKOLOV. B.S.; DOMRACHEV, S.M.; TIKHIY, V.N.; POZNER, V.M., FORSH, N.N.; LYUTKEVICH, Ye.M.; SLAVIN, V.I.; SAZONOV, N.T.; SAZONOVA, I.G.; SHUTSKAYA, Ye.K.; KRASNOV, I.I.; KALENOVA, G.N.; VINOCRADOV, A.P., glav. red.;

[History of the geological development of the Russian Platform and its margins] Istoriia geologicheskogo razvitiia Russkoi platformy i ee obramleniia. Moskva, Nedra, 1964.
251 p. ___[Maps] Karty. 981. (MIRA 18:4)

ZARRINA, Ye.P.; KAPLYANSKAYA, F.A.; KRASNOV, I.I.; MIKHANKOV, Yu.M.; TARNOGRADSKIY, V.D.

Periglacial formation in the West Siberian Plain. Mat. VSEGEI Chet. geol. i geomorf. no.4:54-104 '61.

(MIRA 17:5)

KRASNOV, I. I.

"Regional unified and correlative stratigraphic scheme of the Quaternary of the European part of the USSR."

report submitted for the 7th Intl Cong, Intl Assoc for Quaternary Research, Boulder & Denver, Colorado, 30 Aug-5 Sep 65.

CATESHUM, G.S.; KRASHOV, I.I.

International Map of the Graterrary Sediments of Europe made on a 1:2,500,000 scale. Biul. Ken. chetv. per. no.30:27-57 465. (IMFA 19:2)

13

ACC NR. AP6015763 SOURCE CODE: UR/0048/66 63 AUTHOR: Stoyanov, P. Moseyev, V. V.:

Krasnov. I. V.

ORG: none

TITLE: Magnetic electrostatic deflecting system for an electron microscope illuminating assembly /Report, Fifth All-Union Conference on Electron Microscopy held in Sumy 6-8 July 19657

SOURCE: AN SSSR. Izvestiya. Seriya fizicheskaya, v. 30, no. 5, 1966, 774-777

TOPIC TAGS: electron microscope, electric field, magnetic field, electron optics, prism, aberration

ABSTRACT: The aberrations of electrostatic and magnetic deflecting systems have been investigated experimentally in order to evaluate their possibilities for use as deflecting systems in high resolution electron microscopes. The experiments were performed by deflecting beams of small circular cross section through different angles up to about 3° and recording the cross section shape of the deflected beam. A number of photographs of the deflected beams are presented. Double deflecting systems (deflection of the beam first in one direction and then in the opposite direction) with total deflections up to about 1.50 were tested. The purely electrostatic systems had considerable astigmatism, but when one of the deflectors was a magnetic system with astig matism corrected, as proposed by P.A.Stoyanov and V.V.Moseyev (Radiotekhnika i elek-

Card 1/2

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ACC NR: AP6015763	
4	4
tronika, 8, No. 7, 1169 (1963)) and by P.A.Stoyanov (Izv. AN SSSR. Ser. fiz., 27, 1239	
(1963)), the resultant astigmatism was small and could be corrected in the second condensing lens. Corrected magnetic deflectors were tested at deflection angles up to	1
and slightly beyond 30. The corrected systems showed practically no third order aber-	P
rations, although small fifth order aberrations were perceptible at the largest de-	
flections. The magnetic deflectors showed considerable come when they were mounted	
too close to the iron wall of the housing, but it proved to be possible to correct	
this. It is concluded that a corrected magnetic deflecting system can be employed to	
achieve dark field illumination without significant deterioration of the resolving	
power of the microscope. Orig. art. has: 3 figures.	
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KRASNOV, I. V. M.; STEPANOV, A. V.

Photoelasticity

Optical methods of investigating centers of disintegration. Zhur.eksp.i teor. fiz. 23 no. 2, 1952

Monthly List of Russian Accessions, Library of Congress, December 1952. UNCLASSIFTED

KALASHNIKOV, N.V.; KRASNOV, K.A.

[Automobile filling stations] Avtomobil'nye zapravochnye stantsii. Moskva, Izd-vo Ministerstva kommunal'nogo khoziaistva RSNSR, 1952. 187p. (MLRA 6:8)
(Automobiles--Service stations)

KLEYNERMAN, Yu.A., inzhener; KRASHOV, K.A., redakter; SHELUKHIN, A.S., redakter; KOGAN, F.L., tekhnicheskiy redakter.

[Garage and repair equipment; catalog and manual] Garazhnee i remontnee eberudevanie; katalog-spravechnik. Meskva, Nauchnetekhn. izd-ve avtetransp. lit-ry, 1955. 179 p. (MRA 9:6)

1.Glavnyy inzhener tresta pe preizvedstvu garazhnege eberudevanya (fer Krasnev). (Autemebiles--Repairing) (Service stations)

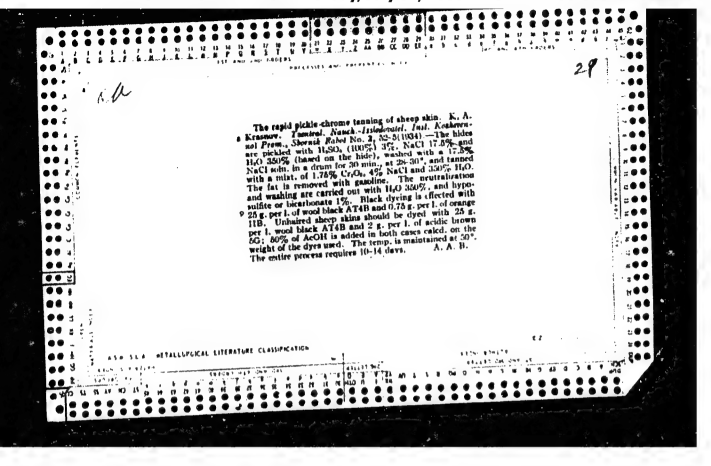
BEREZKIN, Vasiliy Ivanovich; KRASNOV, Konstantin Alekseyevich; MARTENS, S.L., red.; MAL'KOVA, N.V., tekhn.red.

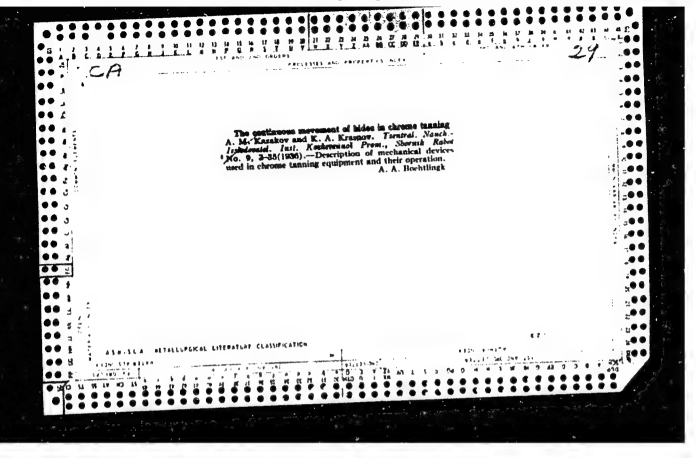
[Equipment for garages and stations servicing automobiles]
Oborudovanie dlia garashei i stantsii obslushivaniia avtomobilei.
Moskva, Nauchno-tekhn.izd-vo M-va avtomobilinogo transp. i
shosseinykh dorog RSFSR, 1959. 273 p.
(Garages--Equipment and supplies)
(Service stations--Equipment and supplies)

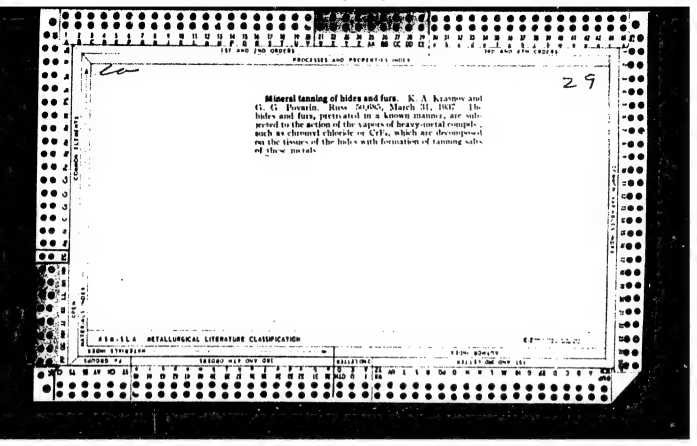
ERECKIN, Vasiliy Ivanovich; MC-TO-V, Monetentin Clekoeyevich;
YABLOKOV, V.1., red.

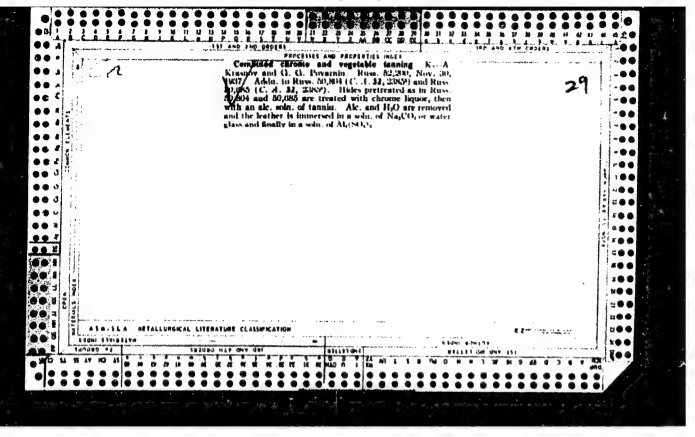
[Equipment for garages and service stations; Oborudovanie dlia garazhei i stantsii obsiuzbivenica avtembliei. 124.2., perer. i dop. Mockva, Transport. 1964. 406.1.

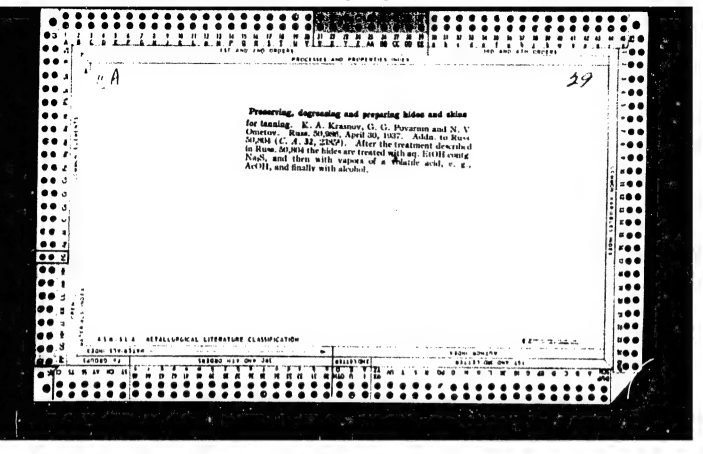
(MIR. 7:7)

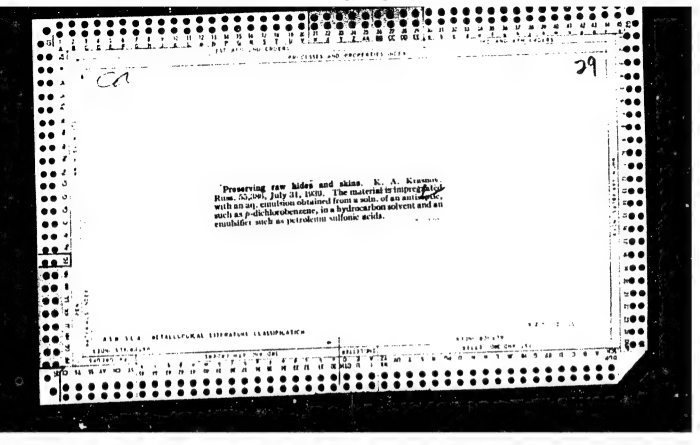






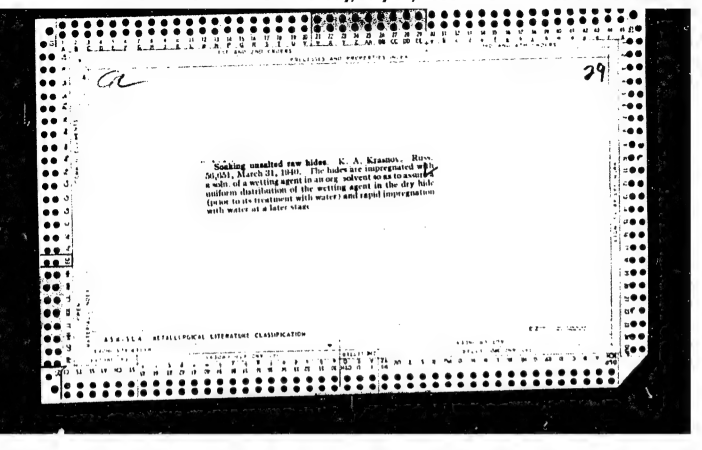


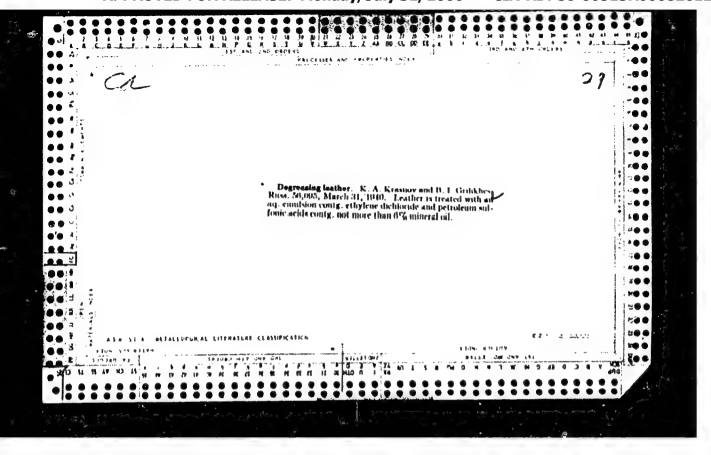




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KRMSMCV, K. A.

Wool Trade and Irdustry

Cleaning the woolly covering off sheared sheepskin. Log. prom. 12 no. 5 (1952)

9. Monthly List of Russian Accessions, Library of Congress, August 1955, Uncl.

"APPROVED FOR RELEASE: Monday, July 31, 2000

CIA-RDP86-00513R000826120

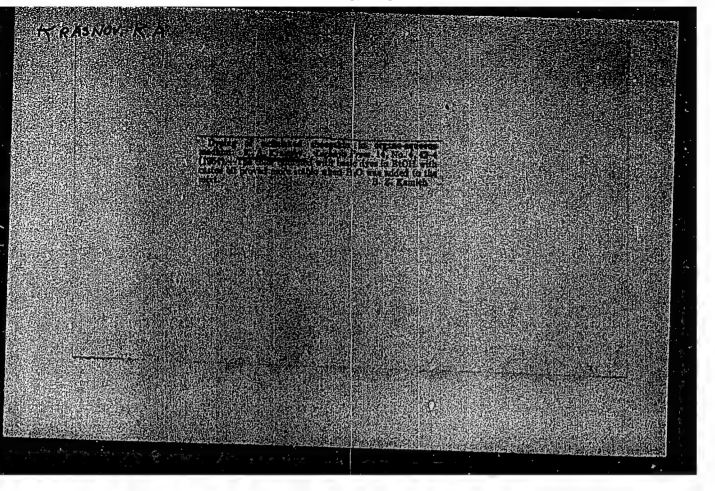
KRASHCV, K. A.

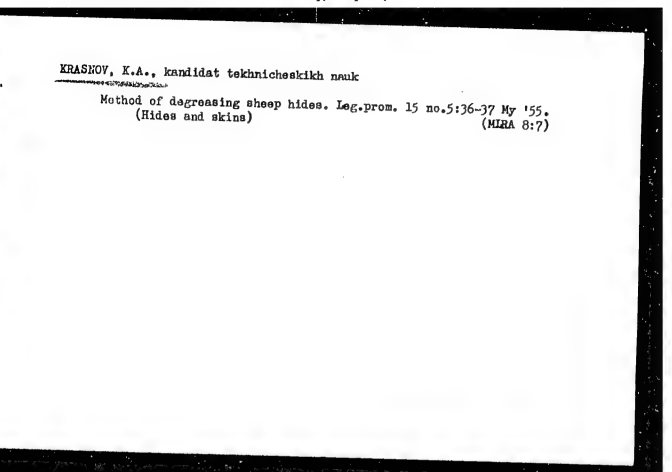
Leather

Improving sheepskins for fleece-lined garments. Leg.prom. 12 No. 6, 1952.

9. Monthly List of Russian Accessions, Library of Congress, Cctober 195%, Uncl.

"APPROVED FOR RELEASE: Monday, July 31, 2000 CIA-RDP86-00513R000826120

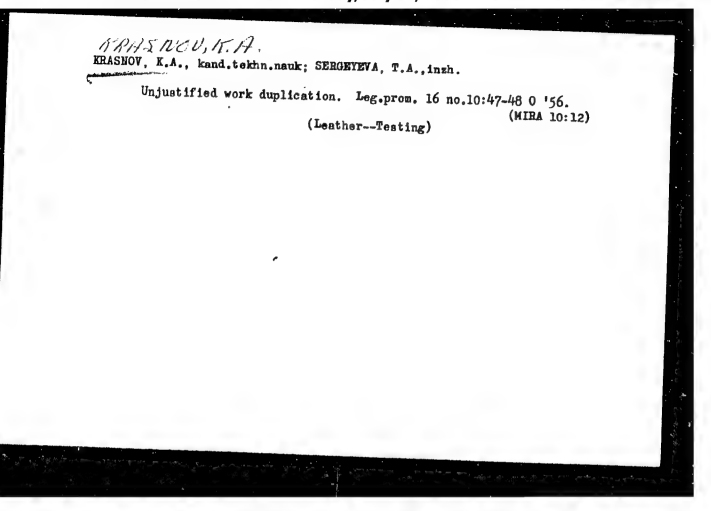




Treatment of fur hides with formaldehyde fumes. Lag.prom. 15 no.9: 29-31 S '55. (MIRA 9:1)

(Hides and skins) (Formaldehyde)

Mays of processing leather and fur. Leg.prom.16 no.2:37-38 P 156.
(Leather industry) (Fur) (MLRA 9:7)

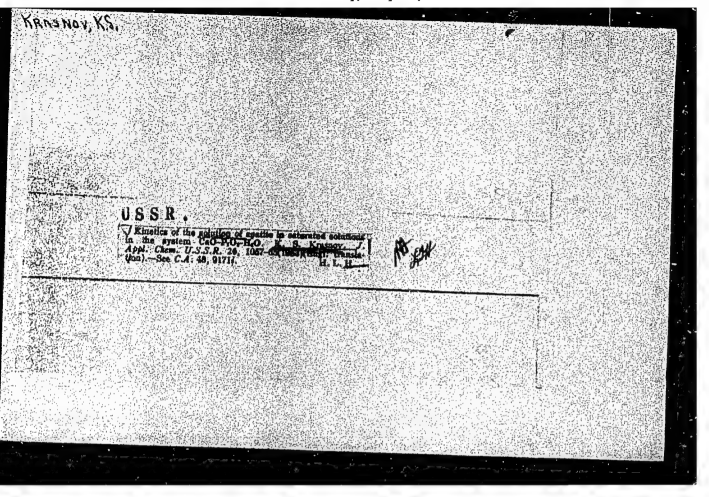


KRASNOV, K.A., otv. za vypusk; YAELOKOV, V.I., red.; BODANOVA, A.P.,

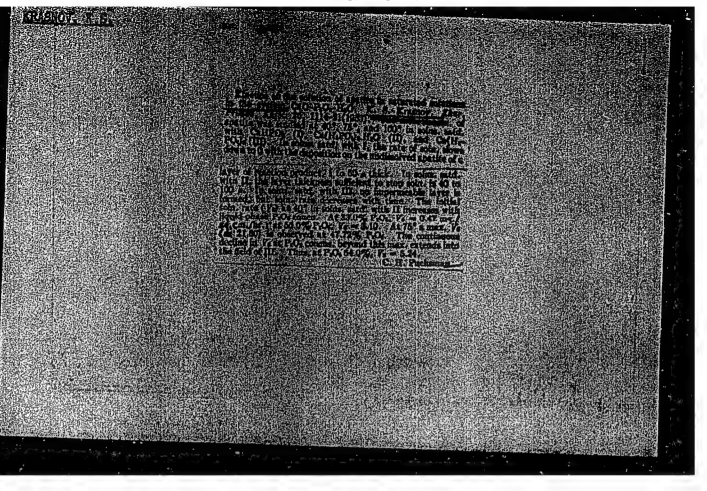
tekhn. red.

[Garage and repair equipment; catalog-handbook]Garazhnoe i remontnoe oborudovanie; katalog-spravochnik. Moskva, Avtotransizdat, 1962. 278 p. (MIRA 16:3)

1. GARO, trust, Moscow. (Motorvehicles-Maintenance and repair) (Garages-Equipment and supplies)



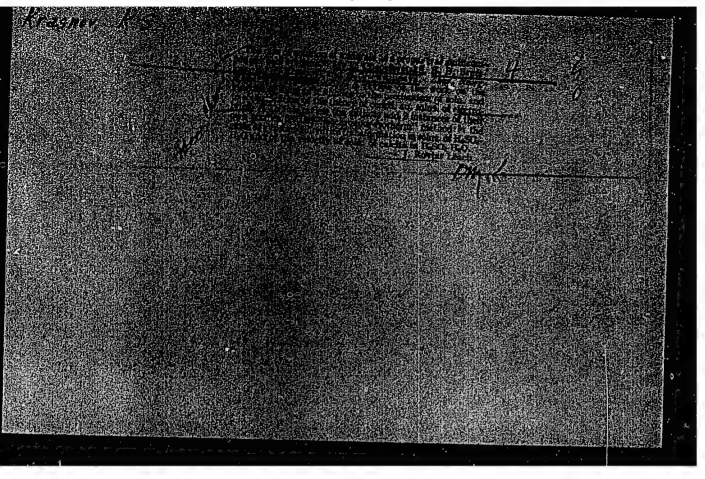
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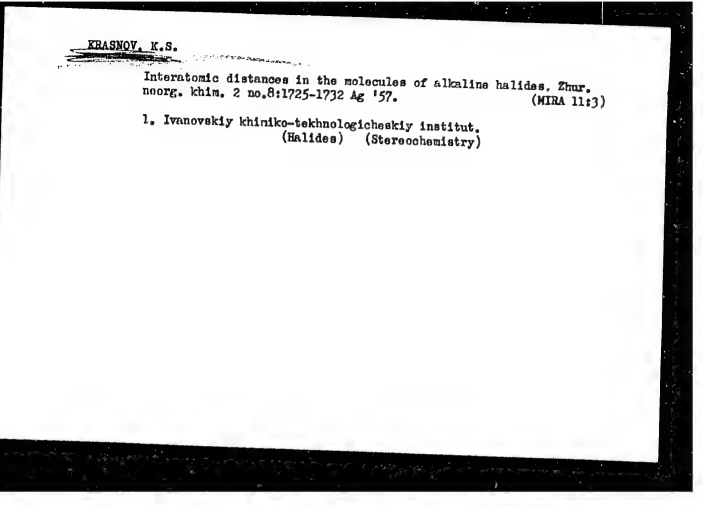


KRASNOV, K.S.

Isotherms of the rate of dissolving apatite in saturated solutions of the ternary system: CaO -- P2O5 -- H2O. Zhur.prikl.khim. 28 no.12:1275-1284 D '55. (MLRA 9:3)

Kafedra khimii Murmanskogo vysshego morekhodnogo uchilishcha.
 (Apatite)





KRASMOV, K.S.

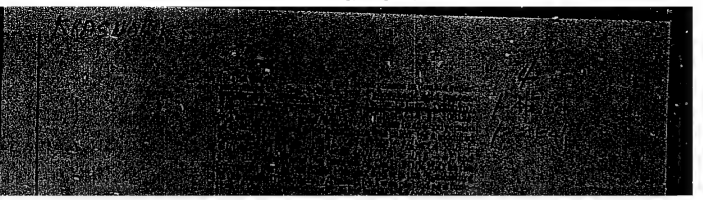
Conditions for producing double superphosphate. Zhur.prikl.khim. 30 no.1:25-32 Ja 157.

(MLRA 10:5)

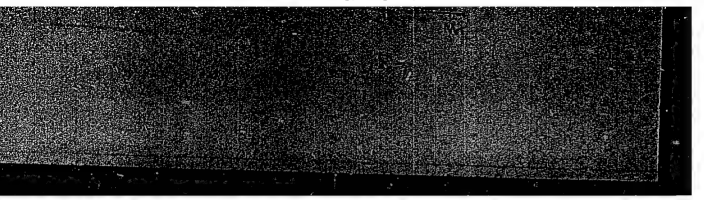
1.Ivanovskiy khimiko-tekhnologicheskiy institut.

(Apatite) (Phosphates) (Systems (Chemistry))

"APPROVED FOR RELEASE: Monday, July 31, 2000 CIA-RDP86-00513R000826120



"APPROVED FOR RELEASE: Monday, July 31, 2000 CIA-RDP86-00513R000826120



5(1) AUTHOR:

Krasnov, K. S.

sov/153-58-3-18/30

TITLE:

Dissolution of Apatite in Phosphoric Acid Partly Neutralized With Magnesium. (Raatvoreniye apatita v fosfornoy kislote, chastichno neytralizovannoy magniyem) The Degree of Neutralization and the Dissolution Velocity (Stepen' neutralization is skerost! restvoreniya)

neytralizatsii i skorost' rastvoreniya)

PERIODICAL:

Izvestiya vysshikh uchebnykh zavedeniy. Khimiya i khimicheskaya tekhnologiya, 1958, Nr 3, pr 100 - 104 (USSR)

ABSTRACT:

The phosphorites of the Kara-Tau contain larger amounts of dolomite. For this reason, magnesium sulfate is formed in the initial stage of sulfuric acid treatment to superphosphate. This magnesium sulfate becomes well soluble magnesium sulfate at a later stage (Ref 1). The liquid superphosphate phase therefore forms a buffer solution consisting of phosphoric acid and its mono-substituted magnesium salt with a considerably higher pH value than that of pure acid. In the next stage this mixture decomposes the rest of the not yet decomposed phosphorite fluoro apatite (or hydroxyl apatite).

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decomposed phosphorite fluoro apatite (or hydroxyl apatite)
This decomposition takes place much more slowly in this
liquid phase than in pure H₂PO₄, and this is the negative

Dissolution of Apatite in Phosphoric Acid Partly Neutralized With Magnesium. The Degree of Neutralization and the Dissolution Velocity 507/153-58-3-18/30

effect of MgCO, in the raw material (Ref 2). For this reason, at the NIUIF (Nauchnyy institut po udobreniyam i insekto-fungisidam im. Samoylova = Scientific Institute for Fertilizers and Insectofungicides imeni Samoylov) and at the laboratory of the author investigations were carried out concerning the problems mentioned in the title. The physical and chemical bases of the acid decomposition of minerals were investigated (Refs 5 - 7), and it was found that it is a typical case of a heterogeneous reaction limited by the diffusion and taking place according to the solution by Shchukarev-Nernst. Assuming that the rate of dissolution is limited towards the apatite surface by the diffusion of the hydrogen ions of the acid, the equation mentioned is transformed into:

$$\mathbf{v} = \begin{bmatrix} \mathbf{k} & \mathbf{H}^{+} \end{bmatrix} \qquad (1),$$

where v denotes the specific rate of dissolution, i.e. the amount of substance that passed into the solution from a surface unit within a unit of time. The equation (1) cannot be used for more concentrated solutions (Ref 3). Therefore

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Dissolution of Apatite in Phosphoric Acid Partly Neutralized With Magnesium. The Degree of Neutralization and the Dissolution Velocity 507/153-58-3-18/30

the author suggested the empirical formula: $v=v_0$. $e^{-\beta Z}$ (2) for the case of the Kara-Tau phosphorites (Ref 3), where v_0 denotes the rate of dissolution in the pure acid with the same P_2O_5 content as the phosphorus magnesium mixture, Z the degree of neutralization of the first H^1 ion of the acid in the mixture and β a constant at the corresponding P_2O_5 content in the solution. The present paper is to prove the usefulness of the equation (2) for various temperatures. This was experimentally carried out at 50 with data of reference 4 being used (at 40 and 75°). The experimental part was carried out by the students L. N. Golubeva and A. I. Mitina. Conclusions: 1.-The exponential dependence of the rate of decomposition of apatite in a partly neutralized phosphoric acid (v) on the degree of neutralization of the acid (Z) was proved. The empiric formula (2) suggested is correct within a wide range of concentration (Z=0-40%) and temperature (25-75°). The coefficient

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Dissolution of Apatite in Phosphoric Acid Partly Neutralized With Magnesium. The Degree of Neutralization and the Dissolution Velocity

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β fluctuates at temperature changes within very narrow limits, and is on the average 0.047.-. There are 2 figures, 2 tables, and 8 Soviet references.

ASSOCIATION:

Ivanovskiy khimiko-tekhnologicheskiy institut (Ivanovo Institute of Chemical Technology) Kafedra fizicheskoy i kolloidnoy khimii (Chair of Physical and Colloid Chemistry)

SUBMITTED:

September 10, 1957

Card 4/4

AUTHORS:

Krasnov, K.S., Antoshkin, V.G.

SOV/78-3-7-5/44

TITLE:

I. The Repulsion Coefficient and the Degree of Ionization of Bonds in Alkali Halides (I. Koeffitsiyent ottalkivaniya i stepenionnogo kharaktera svyazi v galogenidakh shchelochnykh

metallor)

PERIODICAL:

Zhurnal neorganicheskoy khimii, 1958, Vol. 3, Nr 7, pp. 1490-1496

ABSTRACT:

According to the formula developed by Rittner the repulsion coefficient e was determined on the basis of the most recent numarical data between the molecular distances and the number of oscillations in the molecules of the alkali halides. It was shown that the statements made by Rittner concerning the amount of 9 in molecules and orystals are wrong. The repulsion coefficient has the same value for all salts in the molecular and crystalline state. The amount for chlorides, fluorides, bromides, and iodides increases in the direction from chlorides to iodides. For fluorides the mean value of the repulsion coefficient 9 is given as amounting to 0.30 \pm 0.01 %, for chlorides to 0.310 \pm 0.01 %, for brom-

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ides to 0.335 \pm 0.01 Å, and for iodides to 0.365 \pm 0.01 Å.

I. The Repulsion Coefficient and the Degree of Ionization of Bonds in Alkali Halides

307/78-3-7-5/44

The values for the repulsion coefficient ρ and the number of oscillations of the molecules ω_c in alkali halides are calculated. For several alkali salts α_c was corrected. The mean value for the repulsion coefficient of molecules in alkali halides was determined as amounting to 0.33 \pm 0.01 Å. There are 1 figure, 4 tables, and 17 references, 5 of which are Soviet.

SUBMITTED:

Juna 28, 1957

1. Alkali halides—Analysis 2. Alkali halides—Ionization 3. Molecules—Vibrations 4. Mathematics—Applications

Card 2/2

AUTHOR:

Krasnov, K. S.

SOV/78-3-9-1/38

TITLE:

The Relationship Between the Radius of the Non-Deformed Gas Ions and Several Physico-Chemical Characteristics of the Ions (Svyaz' mezhdu radiusom nedeformirovannogo "gazovogo" iona ! nekotorymi fiziko-khimicheskimi kharakteristikami iona)

PERIODICAL:

Zhurnal neorganicheskoy khimii, 1958, Vol 3, Nr 9, pp 1993-1998

(USSR)

ABSTRACT:

In the present paper the interdependence between the gas ion radii on the one hand and the ionization potential and the radius of the crystals of the ions on the other hand was investigated. The relation between the radii of the non-deformed gas ions and the crystallomohemical radii as well as the relation between the radii of the nor-leformed gas ions and the ionization potential were investigated. The results obtained give evidence of a functional interdependence existing between the radii of the ideal non-leformed gas ion r and the physico-

chemical constants of the ion. There is a linear interdependence e. g. between the value $1/z_0$ and the ionization potential,

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between r and the crystallo-chemical radii, and between r and

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The Relationship Between the Radius of the Non-Deformed Gas Ions and Several Physico-Chemical Characteristics of the Ions

other constants that are functionally related to crystallochemical radii. The existing simple functional relation between r_0 and the important physico-chemical constants confirms that r_0 has to be regarded as a physico-chemical constant of the ideal non-deformed gas ions. The radii of the non-deformed gas ion r_{oFr}^+ and the crystallo-chemical radii of this ion were computed and the intra-molecular distances of the halide salts of francium were determined. There are 2 figures, 5 tables, and 17 references,

ASSOCIATION:

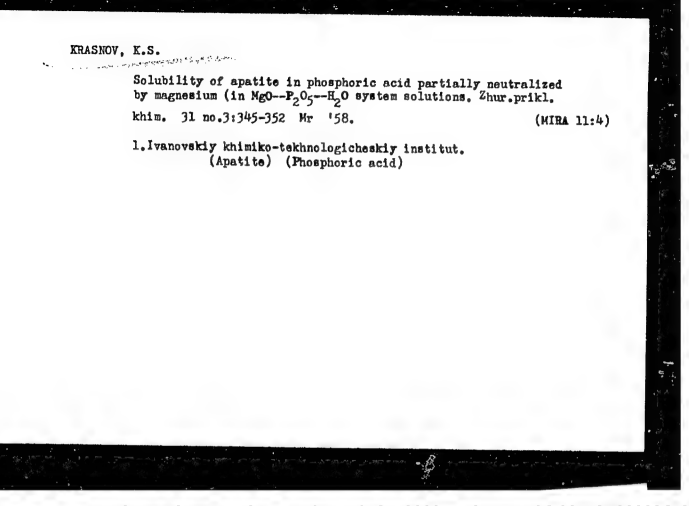
Ivanovskiy khimiko-tekhnologicheskiy institut (Ivanovo Chemical and Technological Instituta)

SUBMITTED:

June 28, 1957

9 of which are Soviet.

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5.2600

TITLE:

Krasnov, K.S.

67034

sov/153-2-5-10/31

AUTHOR:

The Ion Radii and the Nature of the Linkage in Molecules of Metal Selts of the 2nd Group in the Periodic System

Izvestiya vysshikh uchebnykh zavedeniy. Khimiya i khimicheskaya tekhnologiya, 1959, Vol 2, Nr 5, pp 702-705 (USSR) PERIODICAL:

The interatomic distances in gaseous compounds in which the ABSTRACT:

ion character of the linkage prevails can be approximately calculated as the sum of the ion radii in the molecules as it is done in the case of crystals (Refs 1-3). A similar calcula-

tion in more complicated molecules would be of interest. correctness of a constructed ion model of the concerned molecule can be judged by comparison of a calculated and an experimentally determined interatomic distance. For this, of

course, not empirically obtained radii (Ref 3) but as it is done in crystal chemistry, - radii calculated on the basis of theoretical conceptions would be necessary (as in Ref 4). The author attempts to solve this task for the halogen compounds of the 2nd group of the periodic system; there is no uniform

opinion on the nature of their linkage. As a standard for the calculation of the ion radii r+, the author selected inter-

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The Ion Radii and the Nature of the Linkage in Molecules of Metal Salts of the 2nd Group in the Periodic System

atomic distances R in the molecules LiH (Ref 5), NaF (Ref 6), KCl, RbBr, and CsJ (Ref 2) which are built up of isoelectronic ions. For radii of several ions of the latter type, the relation $r_{+} = \frac{C}{2}$ (1) is given in quantum mechanics

where C is the constant of the series, and Z the effective charge of the nucleus (Ref 7). From this, the author obtains for

of the nucleus (Ref 7). From this, isoelectronic ions in the molecule We Hal: $\frac{r_+}{r_-} = \frac{Z_-}{2x}$ (2). After calculating

the effective nuclear charges of the ions in the standard molecules based on the screening constant of L. Pauling (Ref 4), he obtains the values of the halogen radii in the molecules (Table 1, in which for comparison the radii from Refs 2 and 3 are also listed). The ion radii of the halogens were used for the calculation of the radii of the corresponding isoelectronic ions of the metals in the 2nd group (based on (1)) (Table 1). Table 2 lists the electronographically established values of

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The Ion Radii and the Nature of the Linkage in Molecules of Metal Salts of the 2nd Group in the Periodic System

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the interatomic distances (Refs 8,9). They are compared with the values, calculated as the sum of the corresponding ion radii in the molecules. They are in accordance (within the range of the experimental error) with the experimentally calculated distances for salts of Be - Ba. For the compounds of the by-groups, excepting fluorides, there are large deviations (0.2 - 0.3 A). This accordance may serve as one of the criteria to show that in molecules ionic linkage character prevails, especially in the halides of Be - Ba. Any system of ion radii in the molecule which is only based on the additivity principle is incomplete; such a system gives good results as long as the polarization effects in the ionic molecules are close to the same effect in the standard molecules. There are 2 tables and 11 references, 6 of which are Soviet.

ASSOCIATION:

Ivanovskiy khimiko-tekhnologicheskiy institut; Kafedra fizicheskoy khimii (Ivanovo Chemical-technological Institute; Chair of Physical Chemistry)

Card 3/4

67034 The Ion Radii and the Nature of the Linkage in Molecules of Metal Salts of the 2nd Group in the SOV/153-2-5-10/31

SUBMITTED:

Periodic System

June 21, 1958

Card 4/4

· 5(4)
AUTHOR:

Krasnov, K. S.

SOV/78-4-3-7/34

TITLE:

Internuclear Distances in Molecules of the Gaseous Halides of Alkali Metals and the System of Radii of Undeformed Gas Ions (Mezh"yadernyye rasstoyaniya v molekulakh gazoobraznykh galogenidov shchelochnykh metallov i sistema radiusov nedeformirovannykh gazovykh ionov)

PERIODICAL:

Zhurnal neorganicheskoy khimii, 1959, Vol 4, Nr 3, pp 530-534 (USSR)

ABSTRACT:

The interatomic distances in alkali halides were calculated from data on the effective values of the ion radii. The experimentally obtained and calculated interatomic distances in the molecules of the gaseous alkali halides are given in table 3. The deviation from the additivity of the ion radii in this system is due to the mutual polarization of the ions in the molecules and varies between 0.01 and 0.03 Å. Equation (5) has been proposed for calculating the radii of the undeformed gas ions:

Card 1/2

 $d = d_0 - \frac{4(\alpha_1 + \alpha_2)}{(n-1)d_0^2} = r_{0+} + r_{0-} - \frac{4(\alpha_1 + \alpha_2)}{(n-1)d_0^2}$

APPROVED FOR RELEASE: Monday, July 31, 2000 CI

CIA-RDP86-00513R0008261200

SOV/78-4-3-7/34

Internuclear Distances in Molecules of the Gaseous Halides of Alkali Metals and the System of Radii of Undeformed Gas Ions

From the interatomic distance d the radii r_0 of all alkali metal— and halide ions in some molecules can be calculated with the aid of equation (5). The calculated radii of the undeformed gas ions are: Li⁺ = 0.756Å, Na⁺ = 0.980Å, K⁺ = 1,244Å, Rb⁺ = 1.361Å, Cs⁺ = 1.481Å, F⁻ = 1.112Å, Cl⁻ = 1.683Å, Br⁻ = 1.864Å, and J⁻ = 2.119Å. The internuclear distances in the molecules LiCl, LiF, NaF, KF and RbF were calculated with the aid of this system of radii. There are 5 tables and 12 references. 4 of which are Soviet.

ASSOCIATION: Ivanovskiy khimiko-tekhnologicheskiy institut (Ivanovo

Chemical Technological Institute)

SUBMITTED: March 29, 1957

Card 2/2

507/78-4-5-2/46

5(4) AUTHORS:

Shteyn, L. M. Krasnov, K. S.,

TITLE:

Energy in the Molecules of the Halides of Alkali The Bord

Metals

(Energiya svyazi v molekulakh galogenidov shchelochnykh

metallov)

PERIODICAL:

Zhurnal neorganicheskoy khimii, 1959, Vol 4, Nr 5,

pp 963-968 (USSR)

ABSTRACT:

The binding energy W in molecules of the halides of alkali metals was calculated at O according to the formula by

Rittner (Ref 4):

 $W = g' + Ae^{-r/Q} \cdot \frac{c}{-6} + \frac{hv_0}{2}$ (1)

 $\varphi = -\frac{e^2}{\nu} - \frac{e^2(a_1 + a_2)}{2r^4} - \frac{2e^2a_1a_2}{r^4}$

Card 1/4

where $Ae^{-r/\varrho}$ denotes the repulsion energy; $\frac{c}{-6}$

APPROVED FOR RELEASE: Monday, July 31, 2000

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SOV/78-4-5-2/46

The Bond Energy in the Molecules of the Halides of Alkali Metals

of dispersion interaction; $\frac{hy_0}{2}$ - zero point energy; a_1 and a_2 - polarizability of ions;

 $c = \frac{3}{2} \cdot \frac{\mathbf{I}_1 \mathbf{I}_2 \mathbf{a}_1 \mathbf{a}_2}{\mathbf{I}_1 + \mathbf{I}_2}, \quad \mathbf{I}_1 \text{ and } \mathbf{I}_2 \cdots \text{ ionization potentials of the}$ cation and anion respectively. The values \mathbf{W}_1 and \mathbf{W}_2 calculated by means of the formulas (1) and (2) were determined with \mathbf{W}_T . The thermo-chemical values \mathbf{W}_T were calculated according to formula 4: $-\mathbf{W}_T = \mathbf{D}_0 + (\mathbf{I}_0 \cdots \mathbf{E}_0)$ (4)

In formula (4) D_o denotes the dissociation energy of the molecules into atoms in the ground state; I_o — ionization potential of the alkali metals; E_o — affinity of the halides to the electron at 0 K. The data W and W_T concerning fluorides

Card 2/4

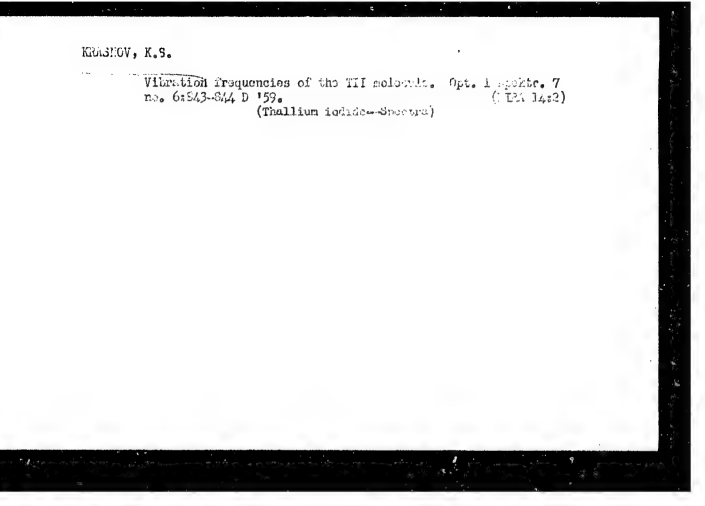
SOV/78-4-5-2/46

The Bond Energy in the Molecules of the Halides of Alkali Metals

are in good agreement. Bond energies of the alkali halides are given by table 1. The differences between theoretical and thermo-chemical values in the binding energy are discussed. The repulsion coefficients for chlorides and fluorides and the following average repulsion coefficients for all alkali halides were determined: NaCl - Q = 0.332; KCl - Q = 0.343; RbCl - Q = 0.355; CsCl - Q = 0.370; NaBr - Q = 0.346; KBr - Q = 0.374 and NaJ - Q = 0.384. The coefficient Q increases from chloride to icdide and from sodium salt to cesium salt. The Q values found are higher than those calculated by Rice and Klemperer (Ref 13). There are 1 table and 13 references, 6 of which are Soviet.

ASSOCIATION: Ivanovskiy khimiko-tekhnologicheskiy institut (Ivanovo Chemical-technological Institute)

Card 3/4



5 (4) SOV /20-128-2-29/59 Krasnov. K. S. AUTHOR: Energy Calculation of Molecules of the Alkaline Earth Halides TITLE: on the Basis of an Ion Model Doklady Akademii nauk SSSR, 1959, Vol 128, Nr 2, pp 326-328 (USSR) PERIODICAL: Under the assumption of ionic linkage a formula for the ABSTRACT: linkage energy in MeX, molecules (Me - alkaline earth, X - halogen) is deduced. The energy, necessary to form a molecule from the cation Me++ and two anions X" with the equilibrium distance Me-X equal ro, if these ions are situated at an infinite distance, is defined as the energy of the molecule MeX2 (T = 0° K). The deduction was made by means of the formula by C. I. F. Böttcher (Ref 5) for electrostatic interaction between two polarizable ions, resulting the formula (5): $U_0 = \frac{-3.5}{r_0} \left(1 - \frac{Q}{r_0}\right) = \frac{4\alpha_2 e^2}{r^4} \left(1 - \frac{4Q}{r_0}\right) + \sum_{i=0}^{h_{i}} \frac{h_{i}}{2} \cdot (9 = 0.358 \text{ Å})$ α_2 = polarization of anion, these values are taken from Landolt- $\frac{h\nu_0}{2}$ is the zero energy, which was Börnstein (Ref 8); Card 1/3

Energy Calculation of Molecules of the Alkaline Earth Halides on the Basis of an Ion Model 501/20-128-2-29/59

neglected in the following calculation). The calculated amounts of \mathbf{U}_0 are cited in table 1 and are compared to measurements by thermochemical means. Further the change $\Delta\mathbf{H}_{298}$ of enthalpy

was calculated on the basis of thermochemical data (Table 1). The differences of the calculated and the thermochemically determined values of U_C vary between -33 kml and zero. The discrepancies amount to an average of %. As all differences are negative a systematic error of calculation is suspected in connection with the determination of the distance Me-X at high temperatures which may be caused perhaps by a covalent component of the linkage. Larger differences may be expected for the same calculation of the molecule HgCl₂ because of stronger covalent linkage. The satisfactory agreement of calculated and thermochemically determined bond energy is therefore not casual but verifies the ionic linkage of MeX₂ molecules. There are 1 table and 10 references, 5 of which are Soviet.

Card 2/3

Energy Calculation of Molecules of the Alkaline Earth Halides on the Basis of an Ion Model SOV/20-128-2-29/59

Earth Halides on the Basis of an 10

ASSOCIATION: Ivanovskiy khimiko-tekhnologicheskiy institut

(Ivarcvo Chemical-technological Institute)

PRESENTED:

May 11, 1959, by A. A. Grinberg, Academician

SUBMITTED:

May 11, 1959

Card 3/3

KRASHOV, K.S.

Radii of ions present in molecules of gaseous inorganic compounds of elements in the main groups of the peroidic system. Zhur. struk. khim. 1 no.2:209-216 Jl-Ag '60. (MIRA 13:9)

1. Ivanovskiy khimiko-tekhnologicheskiy institut.
(Ions) (Chemical bonds)

2h08h \$/186/60/002/006/006/026 A051/A129

21,4200

AUTHOR:

Krasnov, K. S.

TITLE:

Certain thermodynamic characteristics of gasecus francium

halides.

PERIODICAL:

Radiokhimiya, v. 2, no. 6., 1960, 668 - 670

TEXT: The article deals with the computation of certain thermodynamic characteristics of francium halides needed to evaluate the possibilities of separating francium from rabidium and desium by the salt sublimation method. The heat of sublimation is calculated as the difference between the energy of the crystal lattice U and the energy of molecule formation FrG from gaseous ions W (called bond energy for short in the article). The bond energy W₀ at 0°K is calculated on the haris of an ionic model of a molecule according to Pittner's formula (Ref. 11E. S. Rittner, J. Chem. Phys. 19, 1030, 1951):

$$W = \frac{e^2}{r} - \frac{e^2(\alpha_1 + \alpha_2)}{2r^4} - \frac{2e^2\alpha_1\alpha_2}{r^7} - \frac{c}{r^6} + Ae^{-\frac{r}{16}} + \frac{h\omega}{2}$$
 (1)

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24,084

S/186/60/002/006/006/026 A051/A129

Certain thermodynamic characteristics of

where r is the intermolear distance, α_1 and α_2 the polarizability of the ions, C the constant of the dispersion interaction, ω the oscillation frquency, A and ρ constants in the repulsion energy expression. The values of r_e , interatomic distances in FiC molecules, were computed by the author (Ref. 6: K. S. Krasnov, ZhNAh, 3, 9, 1993, 1958). The Varshni formula is suggested for calculating the oscillation frequencies ω_1 ; $r_1 = a + b \cdot k_0^{-\frac{1}{2}}$ (2), where the force constant $k_0 = 4\pi^2/\mu\omega_0$. The values obtained for the oscillation frequencies were found to correspond favorably with those computed by Maksimov (Ref. 7: K. 3. Krasnov, A. I. Maksimov, Opt. 1 spektr. 8, 3, 403, 1960). μ is the true value of the given mass. A comparison of Logs of FrG with that of the Cs, Rb and K halides, showed that Logs of CsF and FrF are lower than those of the corresponding chlorides. This is thought to be the result of the polarization of the voluminous Cs⁺ and Fr⁺ cations by small F⁻ anion. This fact is confirmed by the comparison of the obtained heats of sublimation to the boiling points of cesium and rubidium halides. The values of the thermodynamic functions

 S_{298}^{0} , $-\frac{(z^{0}-H_{0}^{0})}{T}$ and $\frac{(H^{0}-E_{0}^{0})}{T}$ were computed ascording to the

Card 2/3

21,0814 8/186/60/002/006/006/026 AC51/A129

Certain thermodynamic characteristics of ...

statistical thermodynamics formulae. There are 2 tables and 15 references: 8 Soviet-bloc and 7 non-Soviet-bloc. The references to the four most recent English-language publications read as follows: A. Honig, M. Mandel, M. Stitch, C. Townes, Phys. Rev., 96, 3, 629, 1954; J. P. Varshni, Trans Favad. Soc., 53, 2, 132, 1957; Y. P. Varshni, J. Chem. Phys. 28, 6, 1081, 1958; H. O. Fritchard Chem. Rev., 52, 3, 528, 1953.

SUBMITTED:

January 15, 1960

Card 3/3

21,085 8/186/60/002/006/007/026 A051/A129

21.4200

Krasnov, K. Ser Krestov, G. A.

TITLE The volatility of francium compounds

PERIODICAL: Radickhimiya, v. 2, no.6, 1960, 671 - 674

TEXTS The authors calculated the pressure ourves of sublimation for all th- halide compounds offrencium, in order to provide data for establishing the possibilities of sublimation methods. These calculations were based on the high what ility of micro-quartities of francium chloride at 900 - 1,000°K (Ref. 2: Yu. B. Gerlit, F. I. Favlotskaya, S. S. Rodin, Khim. nauka i promyshl. 4, 4, 465, 1959) and facilitated by the calculated series of thermodynamic characteristics of solid and gaveous compounds of franctum. The authors also calculated the sublimation curves of rubifium and desium halides to evaluate the possibilities of the method. The following equation was used for the calculations:

$$\Delta Z_{T}^{0} = \Delta E_{298}^{0} - T \Delta S_{298}^{0} - T \int_{298}^{T} \frac{dT}{dt} \int_{298}^{T} [(c_{p})_{gas} - (c_{p})_{sclin}] dT.$$
 (1)

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CIA-RDP86-00513R0008261200 APPROVED FOR RELEASE: Monday, July 31, 2000

24085 3/186/60/002/006/007/026 A051/A129

The velatility of francium compounds

where ΔZ_{i}^{0} is the change of the importantial in the sublimation process, ΔH_{200}^{0} and ΔS_{200}^{0} the change of the steriari enthalpies and entropies in the same process, C_{n} the in-rmal gapacity at constant pressure. The entropy change in the sublimation process ΔS_{200}^{0} is determined as $(S_{200}^{0})_{gas} = (S_{200}^{0})_{gas}$ and $(S_{200}^{0})_{gas} = (S_{200}^{0})_{gas}$.

The thermal capatity of the solid rubidium and design halides is expressed by the power series equation $C_p = a + bT$. The thermal capacity of the gaseous compounds was calculated from the quantum formula:

$$c_{p} = \frac{7}{2} R + c_{E} \left(\frac{\omega}{T}\right) \tag{2}.$$

where $C_{\rm E}$ ($\frac{h}{L}$) is the thermal capacity of the linear harmonic oscillator. The divides were determined from the infra-rel and micro-wave spectra; for the francium halides they were calculated from the value of the given mass. The sublimation curves within the temperature range of 300 - 1,000°K were calculated from the relation

 $-\Delta Z_{\rm p}^{\rm O} = \rm RT \ lnP \tag{3}.$

Card 2/3

24085

8/186/50/002/006/007/026 A051/A129

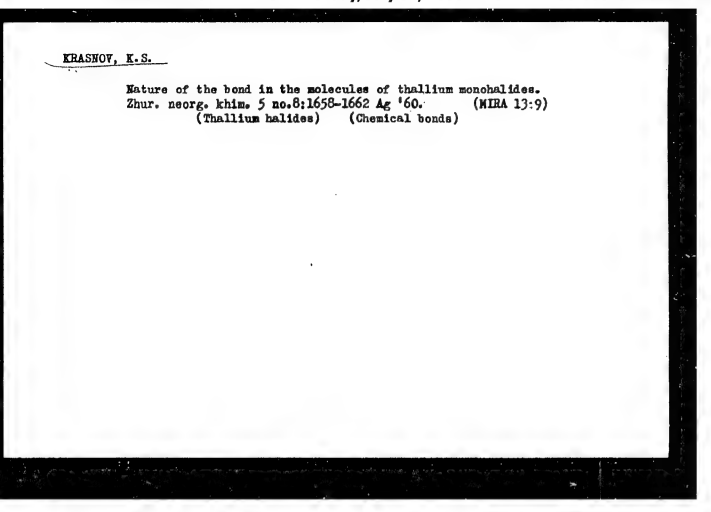
The voiatility of francium compounds

At analysis of these results showed that the sublimation pressure is very low at usual temperatures and increases considerably at high temperatures. The finoride is considered to be the most volatile of all the francium halides and desium compounds. No decomposition of francium indide into elements is expected in sublimation since $\Delta Z \ge 0$ holds true for this process. Indides are more convenient for separation by sublimation than fluorides. This conclusion refers to macro-quantities of substances and cannot be unconditionally applied to micro-quantities of compounds. There are 3 tables and 15 references: 8 Sovietables and 7 non-Sovietables. The references to the English language publications read as follows: A. Honig, M. Mandel, M. Stitch. C. Townes, Phys. Rev., 96, 3, 629, 1954; S. A. Rice, W. Klemperer, J. Chem Phys., 27, 2, 573, 1957; R. F. Barrow, A. D. Count, Proc. Roy. Soc., A,219, 120, 1953; S. A. Rice, W. Klemperer, J. Chem. Phys., 27, 2, 573, 1957; R. F.

SUPMITTED:

February 11, 1960.

Card 5/3

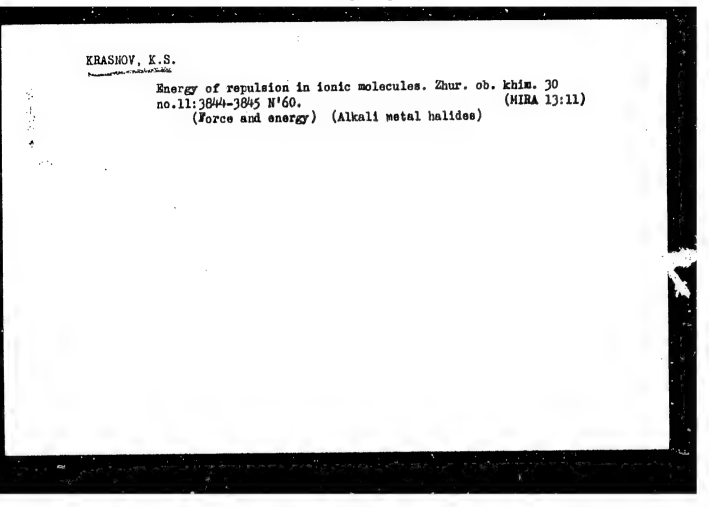


(KRASNOV, K.S.; MAKSIMOV, A.I.

Calculation of the vibration frequencies of diatomic molecules from their reduced mass. Opt. i spektr. 8 no.3:403-406 Mr 160.

(MIRA 14:5)

(Spectrum, Molecular)



21123

S/153/61/004/001/002/009 B110/B203

5.4130 (1273, 1228, 1227)

AUTHOR:

Krasnov, K. S.

TITLE:

Bond energy of halogen compounds of the second group, and

the ion model

PERIODICAL:

Izvestiya vysshikh uchebnykh zavedeniy. Khimiya i khimicheskaya tekhnologiya, v. 4, no. 1, 1961, 38-44

TEXT: For molecules with intermediate character of bond in which ionic interactions prevail, e.g., in TlHal₃, the use of the ion model to calculate the molecular bond energy yields satisfactory results. In this connection, the author had already studied subgroup II A of the periodic system (Ref. 2: K. S. Krasnov, Dokl. AN SSSR, 128, 326 (1959)). The applicability of this calculation method to both subgroups of group II is to be studied here with the use of new data of repulsion coefficients. Let U be the energy change in the formation of MeHal₂ at the equilibrium distance Me - Hal = r from the cation Me²⁺ and two anions Hal infinitely distant. The potential energy of the system MeHal₂ is obtained taking Card 1/8

21123 S/153/61/004/001/002/009 B110/B203

Bond energy of halogen compounds ...

account of: (1) Electrostatic interaction of Me^{2+} and 2 Hal: $W_1 = -2e^2/x - 2e^2/y - 2\alpha e^2/x^4 - 2\alpha e^2/y^4$, where $\alpha = \text{polarizability}$; e = ionic charge; x, y = distances between anions and cation. (2) Electrostatic interaction of Hal between each other: $W_2 = e^2/d - \alpha e^2/\left[d^4(1 + 2\alpha/d^3)\right]$.

(3) Repulsion of the central ion M^{2+} from the dipoles m mutually induced by the anions: $W_3 = \left\{2\alpha e^2/\left[d^2(1 + 2\alpha/d^3)\right]\right\} \cdot (1/x^2 + 1/y^2)$; d = distance of anions from each other. (4) Repulsion of two dipoles m_1 and m_2 induced by the metal ion in the anions: $W_4 = 8\alpha e^2/\left[x^2y^2(1 + 2\alpha/d^3)\right]$.

(5) Repulsion of the completed electron shells of Me^{2+} and Hal: $W_5 = A \exp\left(-x/\rho\right) + A \exp\left(-y/\rho\right)$. (6) Van der Waals dispersing interaction: $W_6 = c/x^6 - c/y^6$; c = London's constant. (7) Difference of values of forward, rotary, and oscillating energies of molecule and free ions:

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21123 s/153/61/004/001/002/009 B110/B203

Bond energy of halogen compounds ...

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Bond energy of halogen compounds ...

absolute zero according to Kirchhoff. Molecules with dependable thermochemical data (Δ H) were chosen for the conversion of sublimation, melting, and evaporation heats. The error of the approximation calculus was below 2-4 kcal. For 10 halides of the main group, the deviation ϵ of the calculated values $\rm U_{o}$ from the thermochemical values $\rm U_{o}$ therm was 3 on an average, and 7% maximum, the degree of ionization of the bond calculated according to Gordi dropping below 65-70%, except for BeJ_2.

A comparison with thallium halides (degree of ionization = 70%, $\varepsilon = 4\%$) suggests, in spite of contrary spectrum interpretations by Western researchers, that bonding ionization prevails up to about 70% and, therefore calculation by the ion model is permissible. The deviations ε are explained as follows: (1) The interatomic distances are measured by electron diffraction at high temperatures. Conversion to 0° K gives smaller and, therefore, more accurate values. (2) It is possible that because of polarization of the gaseous phase, as is the case with alkali halides, also here the real distances measured radiospectroscopically are smaller than those measured by electron diffraction. (3) Influence exerted by the degree of the covalence of the binding. It should be expected in Be

Card 4/8

2]]23 S/153/61/004/001/002/009 B110/B203

Bond energy of halogen compounds ...

compounds in particular. Here, however, the ε -values are small, in BeCl₂ and BeBr₂ even positive. A. Unsöld had already shown that anionic polarization decreases at interatomic distances of nearly 1 A. If values of normal polarizations are used, one obtains good results and a positive deviation for BeCl₂ at r_o = 1.75 A and strong Cl⁻ polarization. The most important components of U_o are Coulombian and polarization interactions as well as quantum-mechanical repulsion. Polarization is between 7% (BaF₂) and 36% (BeJ₂) of the total electrostatic interaction. It increases with decreasing electronegativity difference and is related with many phenomena of covalent binding. Since polarization effect and quantum-mechanical repulsion are nearly equal in chlorides, the simple Coulomb formula U_o = 3.5 e²/r_o gives good results. In the subgroup, only the values for ZnCl_2 and CdCl_2 correspond to the calculated quantities. Considerable deviations (10-20%) are due to mainly covalent binding. This is confirmed by the low equivalent electrical conductivities of the

Card 5/8

21123

Bond energy of halogen compounds ...

S/153/61/004/001/002/009 B110/B203

melts (HgCl₂:BeCl₂ = 1 : 140). In the case of Zn²⁺, Cd²⁺, and Hg²⁺ with 18 electrons, the polarization fraction of the bond energy is larger than indicated in the formula derived. In metal halides with electronegativity below 1.3-1.5 ions, interactions take place, and calculation of energy is possible by the ion model. There are 1 table and 21 references: 10 Soviet-bloc and 11 non-Soviet-bloc. The two references to English-language publications read as follows: R. G. Pearson: J. Chem. Phys., 30, 1537 (1959). A. Büchler, W. Klemperer, J. Chem. Phys., 29, 121 (1958).

ASSOCIATION:

Ivanovskiy khimiko-tekhnologicheskiy institut. Kafedra fizicheskoy i kolloidnoy khimii (Ivanovo Institute of Chemical Technology, Department of Physical and

Colloid Chemistry)

SUBMITTED:

January 19, 1959

W

Card 6/8

KRASNOV, K.S.; MAKSIMOV, A.I.

Use of an ionic model for calculating molecular vibrations of alkaline earth metal halides. Zhur.strukt.khim. 3 no.61703-706 [62. (MIRA 15:12)

1. Ivanovskiy khimiko-tekhnologicheskiy institut.
(Alkaline earth halides) (Molecules--Models)

1956-1251

KRASNOV, K.S.; KASHIRINA, F.D.

Effect os structural factors on the thermodynamic characteristics of basic dye salts. Part 2: Extraction with isoamyl alcohol. Radiokhimiia 4 no.6:638-646 '62. (MIRA 16:1) (Dyes and dyeing) (Isopentyl alcohol)

L 1111;2-61 EWT(1)/EMP(q)/EWT(n)/BDS-AFFTC/ASD-JD/JM ACCESSION HR: AP3000481 S/0153/63/006/001/0167/0170

AUTHOR: Krasnov, K. S.; Svettsov, V. I.

TITLE: Thermodynamic functions of gaseous halides of the alkali earth metals

SOURCE: Izv. VUZ: Miniya i khim. tekhnologiya, v. 6, no. 1, 1963, 167-170

TOPIC TAGS, thermodynamic functions, entropy, enthalpy function, free energy functions, specific heat, vibrational frequencies, molecular force constants, Be halides, Mg halides, Ca halides, Sr halides, Ba halides

APSTRACT: The thermodynamic functions of gaseous alkaline earth halides have not been computed up to the present time because of lack of data on molecular vibrational frequencies. Knowledge of the thermodynamic functions would enable certain calculations to be made in the fields of sublimation and hydration of the halides. The force constants for transverse vibrations, 2k(Delta)/L squared, calculated by the authors according to Pearson's formula, and the frequencies of transverse vibrations, Nu sub 2, calculated to within 50% accuracy according to the valence force-field model, are shown in Table 1 of enclosure 1; included in the table are the longitudinal vibrations, Nu sub 1, and Nu sub 2, as calculated to 3% accuracy by K. S. Krasnov, A. I. Maksimov (Zh. strukturnoy khimii, 3, 707, 1962). Using

Card 1/4

L 11142-63

ACCESSION NR: AP3000481

this information the entropy at 298K, S sup 0, the enthalpy function (H sup 0 - H sup 0 sub 0)/T, and the free energy function -(F sup 0 - H sup 0 sub 0)/T (where F is the Gibbs free energy) at 298, 400, 500, 1000, 1500, 2000, 2500K, were computed for the fluorides, chlorides, broadles, and iodides of Be, Kg, Cs, Er, and Ba, using the model of a rigid rotatorharmonic oscillator. Vibrational frequencies were also used for computing the specific heat of these substances at 298, 500, 700, 1000 1500, 2000, and 2500K. Inaccuracies in the thermodynamic functions introduced by the large uncertainty in Nu sub 2 were found to be insignificant. Orige art. has: 3 tables.

ASSOCIATION: Ivanovskiy khimiko-tekhnologicheskiy institut, Kafedra fizicheskoy i kolloidnoy khimii (Department of Physical and Colloidal Chemistry, Ivanovskiy Chemical Technological Institute)

SUBMITTED: 20Apr 62

DATE ACQD: 21Jun63

ENCL: 02

SUB CODE: CH, PH

NO REF SOV: OCA

OTHER 002

Card 2/4

"APPROVED FOR RELEASE: Monday, July 31, 2000 CIA-RDP86-00513R000826120

KRASNOV, K.S.

Effect of the structural factors on the thermodynamic characteristics of the extraction of basic dye salts. Part 3: Equilibria in the extraction with a "dissociating" solvent. Radiokhimiia 5 no.2: 222-231 '63. (MIRA 16:10)

KRASNOV, K.S.

Ionic radii in molecules of inorganic compounds. Zhur.strukt.khim.
4 no.6:885-891 N-D '63. (MIRA 17:4)

1. Ivanovskiy khimiko-tekhnologicheskiy institut.

Thermodynamic functions of gaseous alkaline earth metal halides.

Izv.vys.ucheb.zav.;khim.i khim.tekh. 6 no.1:167-170 '63.

1. Ivanovskiy khimiko-tekhnologicheskiy institut, kafedra fizicheskoy i kolloidnoy khimii.

(Alkaline earth halides—Thermodynamic properties)

KRASNOV, K.S.; KASHIRINA, F.D.; YATSMIRSKIY, K.B.

Thermodynamics of the extraction of ionic associates as exemplified by the extraction of triphenylmethane dye salts. Trudy Komanal.khim. 14:59-75 163. (MIRA 16:11)

KRASNOV, K.S.; KASHIRINA, F.D.

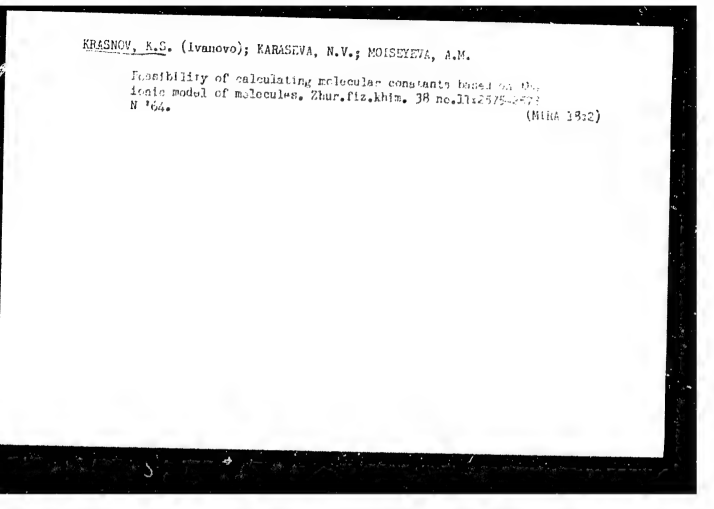
Influence of structural factors on the thermodynamic characteristics of the extraction of basic dye salts. Part 4: Extraction of brilliant green. Radiokhimiia 6 no.2:191 201 '64. (MIRA 17.6)

ZATTERV, A.A.; KRACHOV, K.F.

Some errors in the use of an ionic model of a molecule. Zhur. fiz. khim. 38 no. 3:811-812 Mr 164. (MIRA 17:7)

1. Ivanovskiy khimiko-tekhnologicheskiy institut.

"APPROVED FOR RELEASE: Monday, July 31, 2000 CIA-RDP86-00513R000826120



VCROB'YEV, Nikolay Konstantinovich; GOL'TSSHMIDT, Vladimir Avgustovich [deceased]; KARAPET'YANTS, Mikhail Khristoforovich; KISELEVA, Vera Leonidovna; KRASNOV, Konstantin Solomonovich; LEVINSKIY, Yu.V., red

[Laboratory work in physical chemistry] Praktikum po fizicheskoi khimii. Izd.3., perer. i dop. Moskva, Khimiia, 1964. 383 p. (MIRA 18:4)

KRASNOV, K.S.

Radii of ions in dilute solutions. Zhur. strukt. khim. 5 no.6:834-838 N-D '64. (MIRA 18:4)

1. Ivanovskiy khimiko-tekhnologicheskiy institut.

KRASNOV, K.S.

Binding energy of a AlF₂ molecule. Izv.vys.ucheb.zav.; khim. i khim. tekh. 7 no.2:344-345 *64. (MIRA 18:4)

1. Ivanovskiy khir'ko-tekhnologicheskiy institut, kafedra fizicheskoy i kolloidnoy khimii.

KRASNOV, K.S.; KASHIRINA, F.D.

Effect of structural factors on the thermodynamic characteristics of the extraction of basic dye salts. Part 5: Extraction of salts of methyl green with a two-charge cation. Radiokhimiia 6 no.6:651-655 164. (MIRA 18:2)

KHASNOV, K.S.; KASHIRINA, F.D.

Effect of structural factors on the thermodynamic characteristics of extraction of the salts of basic dyes. Part 6: Composition and structure of solvates in isoamyl alcohol. Radiokhimiia 7 no.2:

(MIRA 18:6)

KRASNOV, K.S.; KARASEVA, N.V.

Revision of spectroscopic dissociation energies for subhalides of group IIa. Opt. 1 spektr. 19 no.1:30-34 Jl "65.

(MIRA 18:8)

KRASNOV, K.S. Energies of dissociation of magnesium and calcium monofluoride compounds and an ionic model of the molecule. Zhur.fiz.khim. 39
(MIRA 18:8) 1. Ivanovskiy khimiko-tekhnologicheskiy institut.

ABASNOV, K.S.; LOBANOV, G.A.; FEOKTISTOV, V.S.

Flectric conductivity of triphenyimethane served tyes in issumyl alcohol. Izv. vys. uchab. zav.; khim. i khim. bakh. 3 no. 4:579-582 *165. (MIRA 18:11)

L. Ivanovskiy khimiko-takhnologicheskiy institut, kafadra fizicheskoy i kolloidnoy khimil.

KRASMOV, K.S.; ZAYTSEV, A.A.

Frequency of stretching vibrations of MeHal3 molecules of lanthanum halides. Zhur.fiz.khim. 39 no.10:2477-2481 0 (MIRA 18:12)

1. Submitted July 15, 1964.

KRASNOV, K.V.

LIVOVSKIY, P.G.; PAL'HOV, Ye.V., professor doktor, retsenzent; KRASHOV, K.V., inzhener, retsenzent; ZAKROCHINSKIY, S.V., inzhener, retsenzent; SHKLOVSKIY, M.B., inzhener, retsenzent; BOGACHEV, I.N., professor doktor tekhnicheskikh nauk, redaktor; AKHUN, A.I., kandidat tekhnicheskikh nauk, redaktor; BARANOV, V.M., kandidat tekhnicheskikh nauk, redaktor; HYZHIKOV, A.A., kandidat tekhnicheskikh nauk, redaktor; CHERNOBROVKIN, V.P., kandidat tekhnicheskikh nauk, redaktor; CHERNOBROVKIN, V.P., kandidat tekhnicheskikh nauk, redaktor; YAKUTOVICH, M.V., kandidat tekhnicheskikh nauk, redaktor; GRISHCHENKO, M.F., inzhener, redaktor; ZASLAVSKIY, I.A., inzhener, redaktor; KROKHALEV, V.Z., inzhener, redaktor; SOSKIH, M.D., inzhener, redaktor.

[Manual for the mechanic in a metallurgical plant] Spravochace ruko-vodstvo mekhanika metallurgicheskege saveda. Isd.3., ispr.i dep.
Moskva, Gos. nauchno-tekhn. isd-vo lit-ry po chernoi i tsvetnoi metallurgii, 1953. 1112 p.

(MLRA 7:4)

(Mechanical engineering--Handbooks, manuals, etc.)

BURDAKOV, D.D.; PANFILOV, M.I.; MEDVEDEV, I.P.; STRUGOVSHCHIKOV, D.P.; NIKOLAYEV, A.M.; KRASNOV, K.Y.

Ways to expand old plants in the Urals. Stal' 16 no.9:818-820 S '56.

(MLRA 9:11)

(Ural Mountain region--Metallurgical plants)

KRASHOV, L.

Therapeutic action of "air vitamins". Okh.truda i sots.strakh.
no.1:52-53 Ja 60. (MIRA 13:5)

I. Chlen zavkoma profsoyuza Dnepropetrovskogo zavoda metallurgicheskogo oborudovaniya.

(Air, Ionized--Therapeutic use)

KRASNOV, L.

Comprehensive plan in action. Okhr. truda i sots. strakh. 5 no.9:8 S '62. (MIRA 16:5)

1. Chlen komissii okhrany truda Dnepropetrovskogo zavoda metallurgicheskogo oborudovaniya. (Dnepropetrovsk--Metallurgical plants--Hygienic aspekts)